Original Article

Role of midkine-progranulin interaction during angiogenesis of hepatocellular carcinoma

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Abstract: Midkine (MK) is a heparin-binding growth factor involved in growth, survival, migration, and differentiation of various target cells and dysregulation of MK signaling is implicated in a variety of inflammatory diseases and cancers. Although MK has been reported to act on endothelial cells and to have proangiogenic effects, the exact role of MK in angiogenesis is poorly defined. Progranulin (PGRN) is a secreted glycoprotein that functions as an important regulator of development, cell cycle progression, cell motility, tumorigenesis, angiogenesis. We screened the PGRN from the hepatic cancer cell cDNA library which was interacted with MK, and confirmed the binding by co-immuno-precipitation and co-localization. During our study, the interaction between MK and PGRN had the important role on the HUVECs proliferation, migration, and tubulogenesis, which indicated the interaction may regulate the angiogenesis, also the in vivo angiogenesis model CAM showed the promotion effect stimulated by MK and PGRN. These findings provide the first evidence linking the association of MK and PGRN and may identify the mechanism of MK during the hepatocellular carcinoma angiogenesis.

Keywords: Midkine, progranulin, protein-protein interaction, hepatocellular carcinoma, angiogenesis

Introduction

Hepatocellular carcinoma (HCC) is one of the most common human tumors worldwide [1]. It is a popular malignancy with more angiogenesis, rapid growth, early metastasis and high mortality. Curative options for HCC include resection and liver transplantation, but prognosis after resection remains unsatisfactory due to a high incidence of postoperative recurrence. HCC is insensitive to chemotherapeutic effects are not satisfactory [2]. Therefore, new therapies with high efficacy, low toxicity, and better specificity for HCC should be considered.

Midkine (MK) is a member of the heparin-binding growth factor family and has been identified as the product of a retinoic acid-responsive gene [3], it is overexpressed in hepatocellular carcinoma and can promote HCC cells’ proliferation and invasion. It also involved in the angiogenesis and tumorigenesis of hepatocellular carcinoma [4], during the previous studied, we found the antisense oligonucleotide (ASODN) that targets MK can suppress the growth of hepatic cancer cells and angiogenesis in nude mice [5, 6]. Additionally, siRNA or an ASODN that targets MK inhibits neointima formation [7] and renal injury after ischemia [8], but the cellular signaling receptors during hepatocarcinoma angiogenesis for MK have not yet been identified and characterized. Study of the molecular basis of MK signal transduction pathways in hepatic cancer cells would further enhance the understanding of their roles in the development and growth of hepatocellular carcinoma.

Methods and materials

Cell lines and culture

Adherent 293T cells (ATCC: CRL-11268, USA) and HepG2 cells (ATCC: HB-8065, USA) were maintained in Dulbecco’s modified Eagle’s medium (Gibco, USA) supplemented with 10% fetal bovine serum (Gibco, USA). Human Umbilical Vein Endothelial cells (HUVECs) (Cascade Biologics: C-003-5C) were cultured in...
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M200 medium supplemented with Low Serum Growth Supplement (LSGS) (Cascade Biologics, USA).

Plasmid constructs

Yeast expression vectors pDBLeu and pEXP-AD502 (both from Invitrogen, USA) are fusion vectors for the linkage of proteins to the Gal4 DNA binding domain and to the Gal4 transactivation domain, respectively. The fragment encoding the full length of MK was amplified by polymerase chain reaction (PCR) and cloned in frame into the SalI/EcoRI sites of pDBLeu (pDBLeu-MK) to serve as bait in the screening assay.

The mammalian expression vector pcDNA3 (Invitrogen, USA) was used to produce the MK and PGRN proteins. The cDNA fragments encoding human full length MK and PGRN was amplified by PCR and subcloned in frame into the HindIII/BamHI sites of pcDNA3 to produce plasmids pcDNA3-MK and pcDNA3-PGRN, which express the proteins of MK and PGRN in the 293T cells.

The recombinant proteins YFP-PGRN and RFP-MK were produced by pEYFP-N1 (Invitrogen, USA) and pDsRed-Monomer-C1 (BD Biosciences Clontech, UK) vectors. The PCR products were subcloned in frame into the Xhol/HindIII sites of pEYFP-N1 and SalI/BamHI sites of pDsRed-Monomer-C1, and the fusion proteins were expressed in the HepG2 cells.

All constructs were verified by nucleic acid sequencing, subsequent analysis was performed using BLAST software (available on the World Wide Web at ncbi.nlm.nih.gov/blast/).

Yeast two-hybrid (Y2H) library screen

Plasmid pDBLeu-MK was used as bait to screen the Y2H human hepatic cancer cells cDNA library according to the PROQUEST™ Two-Hybrid System manufacturer’s protocol (Invitrogen, USA). Briefly, the cDNA library which was inserted into pEXP-AD502 vector and the pDBLeu-MK vector, co-transformed into the yeast MAV203 strain. Yeast cells which contained pDBLeu-MK and pEXP-AD-X cDNA library were laid on SC-Leu-Trp-His+25 mM 3-AT plates. After cultivated at 30°C for 60-72 h, the obvious positive colonies were acquired.

Replica plating were performed by gently pressing the autoclaved filter discs onto SC-Leu-Trp-His plates to transfer the colonies to YPAD plates and selection plates (SC-Leu-Trp plates and SC-Leu-Trp-Ura plates), the former were incubated for X-Gal assay to screen positive colonies, and the latter were for further screening through lacking off Uracil. The fresh colonies of 5 yeast control strains from the glycerol stocks onto SC-Leu-Trp plates were prepared as parallel controls. Finally, the selected plasmids of positive colonies were amplified in E.coli DH5α cells, then plasmids were digested by SalI and NcoI, and the insertion fragmentations were recorded. The positive colonies were sequenced, and the genes were classified according to sequence homology analysis in GenBank (http://blast.ncbi.nlm.nih.gov/). All of the candidate protein coding plasmids and pDBLeu empty vector were co-transformed into yeast cells to identify the self-activity and to re-identify the activity.

Assay of protein-protein interactions using the Y2H system

The full-length MK and PGRN gene were cloned in frame into pDBLeu and pEXP-AD502 respectively. The pDBLeu-MK and pEXP-AD502-PGRN were co-transformed in to MAV203 yeast strains, and then laid on SC-Leu-Trp plates. The co-transformated colonies were then identified on the SC-Leu-Trp-His+25 mM 3-AT plates and analyzed the X-gal activity respectively. The pDBLeu vector+pEXP-AD502-PGRN and the pDBLeu-MK+pEXP-AD502 vector combinations were co-transformated as negative control, while the fresh colonies of 5 yeast control strains were prepared as positive controls.

Assay of protein-protein interactions using the co-immunoprecipitation system

Human 293T cells (5 × 10⁶ cells) were co-transfected with 10 μg of total plasmid DNA using Lipofectamine 2000 (Invitrogen, USA) according to the manufacturer’s instructions. Cells were harvested at 24 h post-transfection and lysed in 1 ml of immunoprecipitation lysis buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1% Triton X-100, 1 mM EDTA) containing 0.1 mM phenylmethylsulfonyl fluoride, 1 μg/ml pepstatin, 1 μg/ml leupeptin, and 1 μg/ml antipain. Extracts were incubated on the ice for 30 min, and the supernatants were collected after 30
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min centrifugation at 4°C (Therom). Antibody (1 μg) was bound to protein A-Sepharose (Santa Cruz, California, CA) by incubation at 4°C for 2 h with rocking. The conjugates were mixed with cell lysis (500 μg of total protein) and incubated overnight at 4°C with rocking. Unbound proteins were removed by extensive washing with lysis buffer. Protein complexes bound to the conjugated antibody were separated by denaturing polyacrylamide gel electrophoresis and subjected to immunoblotting. Anti-MK (N-17) polyclonal antibody (Santa Cruz, CA) and anti-acrogranin (N-19) polyclonal antibody (Santa Cruz, CA) were used for immunoprecipitation, anti-MK monoclonal antibody (Abcam, Cambridge, UK), anti-progranulin monoclonal antibody (R&D system, Minneapolis, MN, CA), anti-rabbit IgG (H+L) HRP antibody (Southern Biotech, CA) and anti-mouse IgG HRP antibody (R&D system, Minneapolis, MN) were used for immunoblotting. Detection was by enhanced chemiluminescence (ECL, Bio-Rad, US).

Co-localization assays of midkine and progranulin

Intracellular co-localization experiments were carried out in HepG2 cells. Cells were grown on collagen-coated cover slips in tissue culture plates to a confluency of 30-40% and were transfected with the plasmids indicated using Lipofectamine 2000 (Invitrogen, USA) according to the manufacturer’s instructions. The expression, localization, and co-localization of the fluorescence-tagged proteins were directly analyzed using the OLYMPUS BX51 fluorescence microscope. Digital images were processed using Adobe Photoshop.

Cell proliferation assay

Human umbilical vein endothelial cells (HUVECs) were plated at a density of 5 × 10⁴/ml into the 96-well culture plates in M200 medium supplement containing 2% (v/v) fetal bovine serum, 1 μg/ml hydrocortisone, 10 ng/ml human epidermal growth factor, 3 ng/ml basic fibroblast growth factor, 10 μg/ml heparin, incubated for 24 h, and subsequently serum-starved for 24 h in serum free M200 medium. HUVECs were either left untreated or challenged with MK, PGRN and MK+PGRN at different concentration respectively (100 ng/ml and 500 ng/ml). As a positive control, 10 ng/ml VEGF-A was added. Cells were incubated for 48 h and 72 h, 10 μl MTT ((3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) (5 mg/ml) reagent was added to the wells and assay was performed after 4 h at 37°C. The absorbance of this colored solution can be quantified by measuring at 490 nm by a spectrophotometer.

Cell migration assay

The migration analysis was performed according to the QCM™ 24-Well Colorimetric Cell Migration Assay manufacturer’s procedure (CHEMICON, MA, USA). HUVECs were serum starved for 24 h. Cells (5 × 10⁵ cells/ml) were then seeded in Boyden chambers (upper chamber), 300 μl per chamber. Lower chambers contained 500 μl of serum free medium or MK and PGRN treatment respectively or co-treatment (500 ng/ml). After 24 h, the cells in the upper chambers were removed, whereas the cells that migrated to the lower chamber were counted after fixation and staining in cell stain solution for 20 min, dipped the insert into a breaker of water several times to rinse, removed non-migratory cells layer from the interior of the insert, lysed the migrated cells in 200 μl of extraction buffer for 15 min at room temperature. Transfer 100 μl of the dye mixture to a 96-cell microtiter plate suitable for colorimetric measurement at 560 nm.

Wound-healing migration assay

Fibrin gel in vitro angiogenesis assay. HUVECs were seeded in the 96-well plates (1 × 10⁴ cells/100 μl), and staved in SFM for 12 h, then the plates were scratched with a thin disposable tip to generate a wound in the cell monolayer. The cells were incubated for additional 24 h in SFM without or with MK and PGRN (20 ng/ml, 100 ng/ml and 500 ng/ml). Cells were analyzed and photographed with a Leica inverted microscope at different time points (0 h, 6 h, 12 h, 24 h).

Tubulogenesis by fibrin gel in vitro angiogenesis assay

The tubulogenesis analysis was performed according to the fibrin gel in vitro angiogenesis assay manufacturer’s procedure (CHEMICON, MA, USA). HUVECs were harvested and counted by standard methods, centrifuged and responded at a final concentration of 5 × 10⁴ cells/ml in SFM without or with MK and PGRN protein (500 mg/ml) reagent was added to the wells and assay was performed after 4 h at 37°C. The absorbance of this colored solution can be quantified by measuring at 490 nm by a spectrophotometer.
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**Figure 1.** Binding of MK and PGRN in yeast. Show is the yeast two-hybrid assay to test the interaction of PGRN which was fused to the Gal4 activation domain and the MK which was fused to the Gal4 DNA-binding domain. Each pair of plasmids, as indicated, were co-transformed into yeast strain MAV203. Yeast transformation were selected on SD-Leu-Trp plates, all of the pairs were growth well on the SD-Leu-Trp plates (A), then tested for growth inhibition on plates containing 25 mM 3-AT (B), the MK/PGRN group and the positive control (strain control E) grow well while the negative control (MK/pEXP-AD502 and pBDLeu/PGRN) did not grow (B). The X-gal activity also have been tested, the MK+PGRN group show the blue colour while the negative control(MK/pEXP-AD502 and pBDLeu/PGRN) did not show blue color (C), the strain A-E were performed as positive control.

ng/ml). Dispensed 30 µl fibrinogen solution into the 96-well plate, gently shook, then added 20 µl per well thrombin to the fibrinogen solution, shook the plate gently immediately, incubated the plate at 37°C for 30 min to polymerize. The prepared cells were seeded onto the fibrin gel (100 µl/well), incubated at 37°C overnight, removed the culture medium completely, added 30 µl per well fibrinogen in each medium to indicated wells of 96-well plat, shook the plate gently, then added 20 µl per well thrombin to the fibrinogen solution, shook the plate gently immediately, and incubated the plate at 37°C for 5 min, added 100 µl per well of SFM without or with treatment. Images of tubular structures were taken after 24 h incubation, using a Leica inverted microscope.

Angiogenesis by chick chorioallantoic membrane (CAM) assay

Angiogenesis analysis in chick chorioallantoic membranes (CAM) was performed as previously described [9]. The fertilized white leghorn chicken eggs were purchased from Wuhu avian breeding center (Huzhou, China), they were incubated under routine conditions (37°C and 70% humidity). On day 6 of development, a square window was opened in the egg shell, after removal of 3.5 ml of albumen allowing detachment of the embryo from the eggshell. The window was sealed with tape, and the eggs were returned to the incubator for 24 h. On day 7 of development, gelatin sponges were cut into a size of 1 mm³ and placed on the top of the CAM under sterile conditions. The sponges were then absorbed with 10 µl of PBS without or with MK and PGRN protein (100 ng/egg), the VEGF (50 ng/egg) was used as the positive control, each group had 8 eggs. CAMs were examined daily until day 10 of development, and then they were fixed in 4% formaldehyde and photographed with a stereomicroscope equipped with a CCD (Pixera EL600, USA). Only CAMs still alive at day 10 were included in this analy-
sis, and then they were all sacrificed by fixing in 4% formaldehyde solution. All of the experiments were carried out according to the standards of animal care as outlined in the Huzhou Central Hospital’s guide for the Care and Use of Laboratory Animals.

**Results**

*Isolation of PGRN as a MK binding partner*

To better understand the biological functions of MK, we performed a yeast two hybrid screen. Briefly, we linked the MK to the Gal4 DNA-binding domain in the plasmid pDBLeu. We used the respective constructs as bait to screen a library of human hepatic cancer cells cDNA expressed as fusion proteins to the Gal4 activation domain in the vector pEXP-AD502. A yeast two hybrid human hepatic cancer cells cDNA library was screened with the construct encoding MK. We screened ~1.0 million colonies and identified 43 colonies that activated with MK gene. Further tests involved the retransformation of yeast with the purified target plasmids and bait. Only 6 of the original 43 yeast clones expressed hybrid proteins that still interacted with the MK bait (Data not show here). One of the positive colonies encoded the growth factor progranulin (accession number NM_002087.2).

*Confirmation of the interaction between PGRN and MK in yeast*

The yeast two hybrid assay was repeated to verify the interaction between the MK and the PGRN. The plasmid encoding the MK linked to the Gal4 DNA-binding domain and the plasmid encoding PGRN fused to the Gal4 activation domain were used to co-transformate into the yeast. Like the yeast control strain A, B, C, D, E as the positive control, our assays indicated that MK interacted with PGRN in yeast, based on the activation of the LacZ reporter gene and the growth phenotypes on SD-Leu-Trp-His+ 25 mM 3-AT plates (Figure 1).

*Binding of PGRN to MK*

The in vivo interaction between MK and PGRN was verified using a co-immunoprecipitation assay in order to determine whether these two proteins are bound in native human 293 T cells. For the co-immunoprecipitation assays, the cell extracts were incubated with either anti-MK antibody or control IgG, and the immunoprecipitated complexes were subjected to a reducing SDS-PAGE and detected with anti-PGRN antibody. A specific PGRN band was present in the immunoprecipitated complexes brought down by anti-MK antibody but not control IgG antibody, demonstrating that PGRN specifically binds to the MK in vivo (Figure 2A). The binding analysis also performed by anti-PGRN antibody...
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Figure 3. MK and PGRN co-localize in the HepG2 cells in vivo. The MK-RFP and the PGRN-GFP fusion protein expression plasmids were co-transfected into the HepG2 cells, the expression of PGRN-GFP (A) and MK-RFP (B) was detected by fluorescence microscopy respectively. The PGRN-GFP and MK-RFP were overlapped in the HepG2 cells (C), the yellow show the overlap areas, the (F) shows the high magnification of overlapped areas. The cells also stained with Hochest33258 to show the nuclear (D, E).

Figure 4. The interaction of MK and PGRN promote the HUVECs proliferation. For in vitro cell growth assay, HUVECs were incubated in SFM and SFM with MK, PGRN, and MK/PGRN proteins (50 ng/ml and 100 ng/ml), the VEGF was used as positive control. The quantification of MTT assay was analyzed at the 24 h, 48 hours and 72 h.

MK and PGRN co-localized in the HepG2 cells

Next, we examined the subcellular localization of MK and PGRN to determine whether these two proteins overlap in the same cell. We first co-transfected human HepG2 cells with plasmids encoding GFP-linked PGRN and RFP-fused MK. As revealed in Figure 3, in the living human HepG2 cells, PGRN is clearly expressed and overlapped with MK. These findings are in agreement with the physical interactions detected in the yeast two-hybrid system and also confirmed by co-immunoprecipitation assays, and suggest that in HepG2 cells, the growth factor MK may be mediated, at least in part, by the PGRN protein.

Interaction of PGRN and MK stimulates the proliferation, migration and tubulogenesis of HUVECs

To elucidate the physiological effect of MK and PGRN interaction in angiogenesis, we used
HUVECs as a model system. We initially performed an in vitro proliferation assay on the 96-well plates. Cells were serum-starved of 24 h and either left untreated or challenged with MK, PGRN, or both growth factors together. As a positive control, cells were stimulated with VEGF (10 ng/ml). We observed almost a three-fold growth induction of HUVECs in the presence of MK and PGRN, respectively.

To investigate whether the interaction of MK and PGRN plays any role in migration of HUVECs, we performed the in vitro “wounding healing” motility assay. The HUVECs were plated at high density in complete medium, after 24 h starvation in serum free medium, confluent HUVECs were wounded and incubated for additional 24 h in serum free medium without or with MK and PGRN protein treatment. In contrast to control, progranulin and midkine both evoked a substantial migration of the cells into the denuded area, and the co-treatment of MK and PGRN indicated a significant promotion of migration, that may suggest the interaction of MK and PGRN have the effectively promote migration of HUVECs. Also, we performed the trans-well migration analysis, the results was consistent with the wound healing experiment.

For the in vitro angiogenesis assay, we cultured the HUVECs under the 3D fibrin matrix, after 24 h incubation in serum free medium without or
with MK and PGRN protein treatment, we found the MK and PGRN can promote the tubulogenesis, and the co-treatment of MK and PGRN have the cooperation effectivity.

Interaction of PGRN and MK promotes the angiogenesis in CAM

The chick chorioallantoic membrane (CAM) assay is a widely used in vivo model that recapitulates the multiple physiological, biochemical and cellular processes that occur during the angiogenic cascade and is relatively rapid for systematically evaluating antiangiogenic agents. Incubation of MK and PGRN on chorioallantoic membranes of 7-day-old chick embryos stimulated an angiogenic response that could be readily photographed after 72 h by stereomicroscopy. MK and PGRN were applied at the dose of 100 ng per egg, the MK and PGRN could stimulate the angiogenesis respectively, but the angiogenesis effect co-induced by MK and PGRN was increased (Figure 7), contrast by the MK and PGRN stimulation respectively, the VEGF was setup as the positive control while the PBS was set as the negative control.

Discussion

Angiogenesis is the process by which new blood vessels are formed [10]. It is a complex multi-step process involving extracellular matrix remodeling, endothelial cell (EC) migration and proliferation, capillary differentiation, and anastomosis [11]. Angiogenesis is observed only transiently during particular circumstances such as reproduction, development, and wound healing. Sustained angiogenesis is characteristic of several pathological conditions including tumor growth [12-15]. Indeed, angiogenesis is essential for tumor growth and it is the interaction of neoplastic with this neovascularature that results in dissemination and metastasis. Endothelial cell proliferation and directed migration are regulated by ploypeptide growth factors and cytokines produced by both the tumor and the host responses to the tumor [16-19]. The variable expression of integrins

Figure 6. The interaction of MK and PGRN effect the tubulogenesis in HUVECs model. For the in vitro angiogenesis analysis, HUVECs were incubated with MK, PGRN, and MK/PGRN proteins (500 ng/ml) according with the Fibrin Gel In Vitro Angiogenesis Assay manufacturer’s procedure (CHEMICON). Images of tubular structures were taken after 24 h incubation, using a Leica inverted microscope.
modulates both the adherence of endothelial cells to components of the basement membrane and the three-dimensional assembly of capillary microvessels [20, 21].

Growth factors play pivotal roles in intercellular communication, and eventually in tissue building and remodeling. The growth factor midkine (MK), also known as neurite growth-promoting factor 2, promoting the growth, survival, migration and differentiation of various cell types [22, 23]. Several reports suggest that MK is a ligand for symdecan-3 [24], anaplastic lymphoma kinase [25], low-density lipoprotein receptor-related protein [26], α4β1- and α6β1-integrin [22], and the receptor-like protein-tyrosine phosphatase-zeta [27]. Clinical reports indicate that MK is overexpressed in a variety of human carcinomas and that high levels of MK in patient serum correlate with poor clinical outcome [28]. Furthermore, high MK expression levels in primary human tumors were shown to correlate with increased tumor angiogenesis [29]. It has also been reported that in
xenografts from two cancer cell lines of epithelial origin, overexpression of the MK gene can cause increased angiogenicity, resulting in enhanced malignant proliferation of cancer cells [30]. Taken together, an increasing number of studies suggest that MK is a candidate molecular target for the therapy for carcinomas. However, the MK’s biologic significance in angiogenesis in general and its mechanism of action in ECs in particular still remains to be clarified.

During our experiments, we use yeast two-hybrid screening to identify protein interaction partners of MK, screening the yeast expression cDNA library using the MK as bait and identified the PGRN, a growth factor that has been previously described in various tumor cells as a direct binding protein of MK.

PGRN is a pluripotent growth factor that mediates cell cycle progression and cell motility, it is a secreted growth factor with high molecular weight that is involved in various biological and pathological processes, including mesothelial differentiation [31], sexual differentiation of the brain [32], macrophage development [33], rheumatoid arthritis and osteoarthritis [34], and wound response and tissue repair [35]. During wound repair it promotes granulation and neovascularization. It is also highly expressed in aggressive cancer cell lines and clinical specimens including breast, ovarian, and renal cancers as well as gliomas [36]. In experimental systems it confers an aggressive phenotype on poorly tumorigenic epithelial cancer cells [37]. Given its actions in wound repair and tumorigenesis PGRN may prove a useful clinical target, both for prognosis and for therapy.

We found that MK and PGRN co-localized in the human Hela cells (Figure 3), and the MK and PGRN complexes could be precipitate by MK or PGRN antibody respectively (Figure 2). These results suggested an in vivo association between MK and PGRN. In line with the previous findings, PGRN interacted with MK in the hepatic cancer cells which was detected by Yeast Two Hybrid system (Figure 1).

Introduced by the the MK alone showed promotion effect on tumor cell growth, while the PGRN also could significantly stimulate tumor cells proliferation. The previously studies found MK and PGRN also can stimulate the endothelial cells proliferation. Our findings show the significant proliferation and migration effect while the endothelial cells were stimulated by MK and PGRN simultaneously (Figures 4, 5). The MK and PGRN also could effect the tubulogenesis, the co-stimulation effect was significantly increased (Figure 6). The in vivo CAM model show the angiogenesis effect was induced by MK and PGRN co-stimulation (Figure 7). These finding suggests that the MK/PGRN complex may play important role during the angiogenesis process. Although the molecular mechanisms underlying the role of MK/PGRN interaction in the angiogenesis remain unclear, it is speculated that PGRN may acts as the co-factor of the MK receptor and may present PGRN to its receptor, followed by the activations of MK-mediated signal transduction and gene regulation pathways.

In conclusion, we have first identified PGRN by yeast two hybrid as a MK-binding protein, and subsequent characterization of this novel association as well as the functional assay showing that the stimulation of endothelial cell proliferation and migration by PGRN growth factors is mediated by MK extend our understanding of the actions of growth factors in angiogenesis and also provide us a potential target for developing and optimizing the therapeutic in tumor angiogenesis and would repair.

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Disclosure of conflict of interest

None.

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MK-PGRN is a growth factor expressed in mesenchymal tissues and is involved in the regulation of angiogenesis. It interacts with angiotensin-converting enzyme (ACE) and is a target for therapies in cancer and other diseases.

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